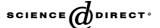


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Prospects for biological control of teasels, *Dipsacus* spp., a new target in the United States

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Abstract

Two closely related teasels (Dipsacales: Dipsacaceae, *Dipsacus* spp.) of European origin have become invasive weeds in the United States. Common teasel (*Dipsacus fullonum* L.) and cutleaf teasel (*Dipsacus laciniatus* L.) have likely been in North America for more than two centuries, having been introduced along with cultivated teasel [*D. sativus* (L.) Honckney], an obsolete crop plant. There are few records of American insects or pathogens attacking *Dipsacus* spp. Invasive teasels have recently begun to spread rapidly throughout much of their current range, for reasons that are not yet known. Common and/or cut-leaf teasel have been listed as noxious in five US states and as invasive in 12 other states and four national parks. Because the family Dipsacaceae is an exclusively Old World family, classical biological control is an important component of the overall management strategy of this weed in the US. Field surveys for natural enemies of *D. fullonum* and *D. laciniatus* in their native ranges and literature reviews of natural enemies of plants in the family Dipsacaceae have yielded 102 species of insects in six orders, as well as 27 fungi from 10 orders, three mites, one nematode, and two viruses. Due to the biennial nature of these weeds, a strategy to assign highest priority to biological control candidates attacking first-year (rosette) plants has been established. Candidates selected for further study based on this strategy include *Chromatomyia ramosa* (Hendel) (Diptera: Agromyzidae), *Longitarsus strigicollis* Wollaston (Coleoptera: Chrysomelidae), *Epitrimerus knautiae* Liro (Acarina: Eriophyiidae), *Euphydryas desfontainii* (Godart) (Lepidoptera: Nymphalidae), *Erysiphe knautiae* Duby (Erysiphales: Erysiphaceae), and *Sphaerotheca dipsacearum* (Tul. and C. Tul.) (Erysiphales: Erysiphaceae).

Keywords: Classical biological control; Invasive species; Dipsacus fullonum; Common teasel; Dipsacus laciniatus; Cutleaf teasel; Dipsacus sativus; Cutlivated teasel; Chromatomyia ramosa; Longitarsus strigicollis, Epitrimerus knautiae; Erysiphe knautiae; Euphydryas desfontainii; Sphaerotheca dipsacearum

1. Introduction

Teasels (*Dipsacus* spp.; Dipsacales: Dipsacacae) are increasing their status as invasive alien weeds in non-agricultural habitats in the US (Sforza, 2004). Invasive teasels occur in 43 US states, being absent only from the extreme southeastern states, North Dakota, Alaska, and Hawai'i (Singhurst and Holmes, 2001; USDA, 2004; Rector, unpublished data). Teasels also occur in the Canadian provinces

of Ontario, Quebec, British Columbia (Werner, 1975a), and Manitoba (Environment Canada, 2003). Four states in the western and midwestern US (CO, IA, MO, and NM) have declared *Dipsacus fullonum* L. (common teasel) noxious, and *Dipsacus laciniatus* L. (cutleaf teasel) is considered noxious in Colorado and Oregon. Cultivated teasel, *Dipsacus sativus* (L.) Honckney, is also present in the US. Teasels are listed as invasive by 12 other states and are listed as affecting natural areas in four national parks (USDI-NPS, 2003). This combined status led to the initiation of a government-sponsored biological control program against these species.

The Dipsacaceae sensu lato is an exclusively Old World family, except in cases where species have been moved by

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humans. Thus, no members of the Dipsacaceae are native to the New World (Sforza, 2004). In addition, there are no plants of significant economic importance within the family Dipsacaceae (Bailey, 1951). The center of origin of the subgenus *Dipsacus* L., which includes all invasive *Dipsacus* spp. in N. America, appears to be in southern Europe, due to the greatest diversity and greatest number of endemic species in that region (Fig. 2A) (Verlaque, 1985). A molecular genetic study is underway to investigate the centers of origin of *D. fullonum* and *D. laciniatus* and the geographical origins of genotypes of these species that are invasive in the US.

This paper documents the known herbivores and pathogens of plants in the family Dipsacaceae in general, and those of *D. fullonum* and *D. laciniatus* in particular. The findings of initial field surveys in the native ranges of *D. fullonum* and *D. laciniatus* are presented, as well as a summary of the existing literature and database resources. Using this information, we make a case for the prioritization of those herbivores and pathogens that have sufficient potential as biological control candidates to warrant intensive impact and host-specificity testing.

2. Teasels in the United States

2.1. Synonomy and history of the target in the US

There has been some confusion over the synonomy of teasel species. Common teasel (sometimes referred to colloquially as "Indian teasel") has frequently been called D. sylvestris (Huds.) rather than D. fullonum, particularly in the North American literature (e.g., Glass, 1991; Huenneke and Thomson, 1995; Judd, 1983). In addition, those who refer to common teasel as D. sylvestris have sometimes used D. fullonum as the name for cultivated (or "Fuller's") teasel, which is otherwise known as D. sativus. A detailed discussion of this taxonomic issue by Ferguson and Brizicky (1965) concluded that the most appropriate name for common teasel is D. fullonum. In addition, because D. fullonum is the type species of the genus, the species name cannot be changed (Bobrov, 1957). The Weed Science Society of America refers to common teasel as D. fullonum, cutleaf teasel as D. laciniatus, and cultivated teasel as D. sativus (WSSA, 2005), and we will use this nomenclature for the remainder of the paper.

Cultivated teasel heads were grown in the pre-industrial era for use in carding or "teasing" wool fibers (Ryder, 1998). The intentional cultivation of teasel has been documented to as far back as 12th century France (Andrieu-Ponel et al., 2000) and may date to the Roman Empire (Ryder, 1998). Cultivated teasel (*D. sativus*) was still an important crop in Europe during the height of European colonization of other continents. This was likely the species of "fuller's teasel" that was introduced by John Bartram into Pennsylvania in 1728 (Tabor, 2003).

Dipsacus sativus very closely resembles D. fullonum and has long been considered to be domesticated from that species (Darwin, 1859). D. laciniatus is also similar in appear-

ance to *D. sativus*, particularly the seeds and seedheads. Introduction and spread of *D. fullonum* and *D. laciniatus* in N. America (as well as other former European colonies) almost certainly arose, at least in part, from contamination of *D. sativus* seed, although the introductions themselves do not appear to have been recorded in the literature.

Despite its utility in the processing of wool, teasel was never a major crop. Relatively little acreage was needed to fulfill the demands of the industry. For example, in 1920 the entire British demand of 10,000,000 teasels could be produced on less than 400 ha of land (Ryder, 1998). As a result, there is little scientific literature concerning teasel production or its associated pests.

Stoner (1951) described an aphid-transmitted virus disease from a "commercial planting of fuller's teasel ... south of Sunnyvale, Calif.," in May, 1948. Thus, *D. sativus* was still under cultivation in the US in the mid-20th century. Topham (1968) also states that teasels were being cultivated in the states of Oregon and New York in that period. Based on the above dates and locations, *D. fullonum* and *D. laciniatus* have had many opportunities for introduction into and spread across America over the course of two centuries.

Spread of invasive teasels through commerce and general interest in the plant continues. Gardeners plant teasel for its striking appearance and purple flowers, its use in dried flower arrangements, and its attractiveness to butterflies, bumblebees, and natural enemies of crop pests (Judd, 1983). Teasel seed, as well as dried teasel flower arrangements (including seed heads that may to contain viable seed), can be purchased through the internet. Also on the internet, one can find numerous teasel-related homeopathic medicinal items and testimonials to their purported efficacy (e.g., Hall and Wood, 2001; Nature's Health Co, 2001; Teeguarden, 2004). Consumers wishing to utilize the plant for any of these purposes may be contributing to the spread of teasel by growing the plants in their gardens or inadvertently spreading viable seed. Teasel seed has also been used in birdseed mixes and may have spread through commercial birdseed sales.

2.2. Target life history and factors affecting weediness

Common, cultivated and cutleaf teasels are often considered biennials because sufficient energy for reproduction is not gained in the first full year of growth, with bolting and flowering normally occurring in the second year. However, under adverse biotic or abiotic conditions (including herbivory or other natural enemy attack) the plant may need additional years to bolt, becoming less likely to do so with each passing year (Werner, 1975b). Given that reproduction only occurs once, no matter the length of the preceding vegetative period, these three *Dipsacus* species are properly referred to as monocarpic, short-lived perennials.

Seeds of common teasel germinate from spring to late summer (Werner, 1975a), after which rosette leaves and a taproot form. The plant grows vegetatively as a rosette, storing energy in the taproot until there is sufficient storage to sustain bolting, flowering, and seed production; bolting has been linked to rosettes exceeding a minimum diameter of 30 cm (Werner, 1975b). Plants that achieve this size late in the year bolt the following spring. At the terminus of each stem a single ovoid to cylindrical seedhead forms. The seedhead on the central stem is the largest on the plant and it flowers first, usually in midsummer. Seedheads on secondary stems flower after the central head, over the course of up to 40 days (Chuko and Hanyu, 1990).

Although self-pollination appears possible, allogamous fertilization, following cross-pollination facilitated by bumblebees, macrolepidoptera, and other insects, is the most common method of reproduction for D. fullonum (Werner, 1975a) and D. laciniatus (Verlague, 1985). Seeds mature within the head in autumn and most fall from the head before the onset of winter, although some viable seed remain in the head into the following spring (B. Rector, unpublished data). In studies on D. fullonum, Werner (1975c) reported that virtually all of the seeds from a given plant (99.9%) fall within 1.5 m of the plant. Long range seed dispersal occurs mainly due to floating seeds in floodwaters or in other flowing waters (e.g., ditches or streams). Common teasel seeds can float up to 22 days without significant reduction in viability (Werner, 1975a). In years without flooding, dense teasel populations can build up as entire seed loads are successively dumped in one area.

Individual teasel plants compete for resources with neighboring plants by spreading large rosette leaves that shade the ground. Common teasel's taproot can extend deeper than the roots of many of its annual and biennial competitors in North America (Werner, 1975a). A teasel plant can produce up to 40 seedheads, the largest of which can produce up to 2000 seeds. Common teasel has been shown to tolerate elevated salinity levels in comparison to other roadside plant species, thus conferring a competitive advantage to teasel in areas where roads are salted in the winter months (Beaton and Dudley, 2004).

Common teasel occurs in dry-mesic and mesic savannas, wetlands, lake borders, agricultural fields, pastureland, successional fields, and developed land (Iverson et al., 1999). The plant grows best in full sun and in poorly drained soils, especially in areas prone to flooding. It is often found in moderately disturbed habitats, such as along roadsides or in waste areas, where seed germination has been shown to be enhanced (Roberts, 1986). Teasel can colonize prairie and savanna habitats, sometimes resulting in monocultures and the exclusion of native species (Glass, 1991; Huenneke and Thomson, 1995). No one has attempted to calculate an economic value for the impact of these invasions.

3. Management options

Current management options for invasive teasels include herbicide treatment of rosettes, mowing of bolted and flowering stems, or, in environmentally sensitive settings, recruiting volunteers to dig up the deep taproots of rosettes and cut and remove stems of bolted plants (Glass, 1991). Fire is inappropriate where teasel populations occur along roads with heavy vehicular traffic and also inappropriate in many natural settings where the risk of wildfire is important. In addition, the plant's most common habitats are characterized by moist soils, which could obviate the use of fire as a general control practice. Effectiveness of other control methods varies considerably (Cheesman, 1998). Mowing of early season stems is considered ineffective since plants can often bolt a second time, necessitating a second mowing. Mowing flowering plants is only effective if the heads are collected and removed—otherwise the mower can scatter seedheads containing viable seed, even when heads are cut before seed reach full maturity (Cheesman, 1998; Solecki, 1989). Glyphosate and 2–4 D have proven effective in killing teasel, but applications over several years are required to manage an established population (Skolnik, 1999), and their use may be restricted in environmentally sensitive areas, such as near waterways.

Given the difficulties controlling established teasel populations, alternative approaches to control are warranted. Biological control of teasel represents one available option. Because of the species-specific nature of biological control, where candidate agents are chosen after extensive host-specificity testing, it can be an effective option while minimizing effects on non-target species (Quimby et al., 2003). Biological control is a particularly attractive option for teasels in North America because of the close phylogenetic relationship between the two invasive teasel species, *D. fullonum* and *D. laciniatus*, as well as the absence of any economically important or native American members of the family Dipsacaceae.

4. Identification of herbivores and pathogens associated with teasels

Field surveys for herbivores and pathogens that attack Dipsacus spp. have been conducted so far in Bulgaria, France, Greece, Hungary, Italy, Romania, Russia, Slovenia, Spain, Turkey, and Ukraine, from 2001–2004 (Fig. 1). In general, field surveys were conducted by traveling within the known ranges of D. fullonum and D. laciniatus (see Fig. 2A) and stopping to inspect plants for herbivore or pathogen damage whenever these species were encountered. A priori information regarding known locations of either of these species or their natural enemies was used whenever it was available. These surveys (Table 1), as well as literature searches (Table 2), have yielded a large number of natural enemies to screen for their suitability as biological control agents of invasive teasels. To date, the total pool of biological control candidates (BCCs) identified from the combined field and literature surveys includes 102 species of insects in six orders, as well as 27 fungi from 10 orders, three mites, one nematode, and two viruses (see Tables 1 and 2). Much of the natural range of D. fullonum and D. laciniatus, as well as the congeners of these species, remains to be surveyed, including much of northern Europe and Scandinavia, the British Isles, the remainder of eastern

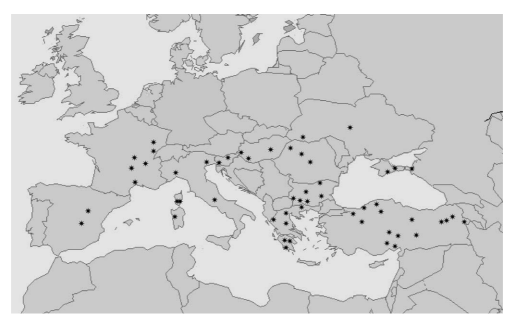


Fig. 1. Survey sites for herbivores and pathogens of Dipsacus spp.

Europe and European Russia, mediterranean northwest Africa, western Turkey and parts of southwest Asia (Fig. 2) (Verlaque, 1985).

In the literature survey, the pool of BCCs was expanded to the broadest logical sense, including those herbivores or pathogens that have been described attacking any plants in the entire family Dipsacaceae (Table 2). An attempt was made to find any reference to *Dipsacus* using all literature searching abstract journals and databases, in the biological, zoological, and agricultural sciences, particular those references pertaining to natural enemies of Dipsacus spp. or any other plant in the Dipsacaceae. Natural enemies found to feed exclusively within this family would not be expected to attack any North American natives or any economically important species. Indeed, a BCC whose primary host is in the Dipsacaceae but is not a *Dipsacus* sp., could become a promising candidate (Hokkanen and Pimentel, 1984), although the first step in testing any such candidate would be to establish that it will feed on the target. Apart from those herbivores that feed on Dipsacaceae, a list of some pollen and nectar feeding insects and other pollinators associated with D. fullonum in eastern N. America is also available (Judd, 1983).

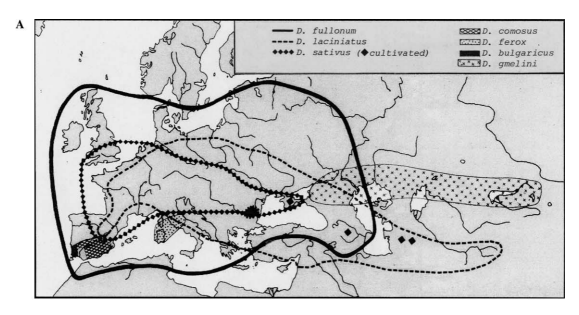
4.1. Prioritzing biological control candidates

In prioritizing intensive research on selected BCCs, a key criterion that must be met is the specificity of the BCC for the target plant. Information on each species's host range in Tables 1 and 2 was sought in the literature and in online databases. An index of specificity was generated using this information to express host ranges relative to the genus *Dipsacus* L. and the family Dipsacaceae (see Tables 1 and 2). Of the 135 BCCs in Tables 1 and 2, 100 are known to attack plants outside the Dipsacaceae. These species, especially those that are highly polyphagous or known to attack

economically important plants, immediately assume a very low priority relative to the other BCCs limited to hosts in the Dipsacaceae. Thirty-two of the BCCs listed in Tables 1 and 2 are only known from hosts in the Dipsacaceae while three have host ranges that are not yet known. Of the 32 BCCs only known from Dipsacaceous hosts, 18 are known to attack *Dipsacus* spp.; and of these 18, five are known only from *Dipsacus* hosts (Tables 1 and 2).

Based on the biennial development of *D. fullonum* and *D.* laciniatus, the requirement of a minimum rosette diameter for successful bolting in D. fullonum (Werner, 1975b), and the diminishing probability of bolting with age (Werner, 1975b), BCCs that attack the taproot or the rosette hold the greatest promise for biological control of teasels. Damage by such agents should either directly or indirectly (in the cases of root- or rosette-feeders, respectively) reduce the storage reserves necessary for bolting and could result in stunted flowering plants with reduced numbers of seedheads or even death without flowering. In addition, damage to the root can open the plant to infection by soil-borne pathogens that would not normally infest a healthy root. Bolting plants and seeds would be important secondary targets. Among the 32 BCCs that are currently only known from hosts in the Dipsacaceae (see Tables 1 and 2), six are known to attack either the roots or rosettes. For many foliage-attacking species, information is not available that indicates whether these specialize on the rosette stage. Five of the 32 BCCs specific to the Dipsacaceae are described from either flowers or seeds.

Other criteria commonly used for prioritizing intensive study of BCCs include but are not limited to severity of damage to the target, coincidence of target and BCC phenologies, and BCC fecundity and distribution (Blossey, 1995; Harris, 1973, 1991; Wapshere, 1974). However, the importance of particular criteria can vary depending on the target, the similarity or difference between native and



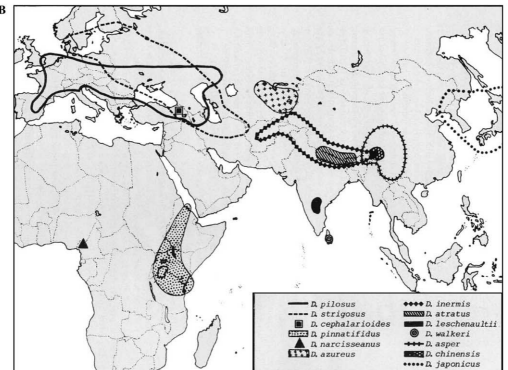


Fig. 2. Native range of the genus Dipsacus L. including the subgroups Dipsacus (A) and Sphaerodipsacus (B) (Verlaque, 1985).

invaded environments, or extreme strengths or weaknesses among members of a given pool of BCCs for one or more of the criteria. From among those candidates listed in Tables 1 and 2, the following BCCs have been assigned the highest priority for going forward into host-specificity and impact testing, based on the above guidelines.

4.2. Candidates already in the US

Thus far, literature and field surveys have yielded evidence of only one native North American, herbivore on teasels. However, this evidence is somewhat anecdotal

and has yet to be confirmed by the authors in the field. According to Covell (1984) larvae of the noctuid moth *Papaipema arctivorens* Hampton feed in rhizomes of thistle, teasel, and burdock in eastern N. America, although the effect that this feeding might have on the teasel plant or on invasive teasel populations is not discussed. Since a native American BCC would have the potential of avoiding many of the quarantine-related permits and testing that foreign BCCs require, the authors will attempt to locate wild populations of *P. arctivorens* in N. America to confirm this insect's host-range as reported by Covell (1984). Preparation of such a BCC for distribution would

Table 1 Identified invertebrate herbivores and pathogens collected by the authors from *Dipsacus* spp. in Europe and Asia Minor in the years 2001–2004

Order	Family	Genus	Species (author)	Damage	Specificity ^a	Range ^c	Reference(s)
Fungi							
Helotiales	Sclerotiniaceae	Sclerotinia	sclerotiorum (Lib.) de Bary	Seedling, bolted plant	X	Co	Brosten and Sands (1986); Cother et al. (1996)
Moniliales	Moniliaceae	Alternaria	alternata (Fr.) Keissl.	Seedling/rosette	X	Co	Widmer, unpublished data
Sphaeropsidales	Sphaeropsidaceae	Phoma	exigua Sacc.	Crown and roots	X	Co	Widmer, unpublished data
Mites Acarina	Tetranychidae	Tetranychus	urticae Koch	Foliage, stems	X	Co	Rector and Sforza, unpublished data
<i>Insects</i> Coleoptera	Cerambycidae	Agapanthia	osmanlis Reiche	Stalk, bolting	G	Eu	Kovacs (1998); M. Rejzek,
	Chrysomelidae	Altica	oleracea (L.)	plants foliage	X	Pa	pers. comm. Warchalowski (2003)
	,	Chaetocnema	concinna (Marsham)	Roots, foliage	X	Eu, As	Doguet (1994); Cagán et al. (2000); Warchalowski (2003)
		Chaetocnema	tibialis (Illiger)	Roots, foliage	X	Med	Doguet (1994)
		Crepidodera	aurata (Marsham)	Foliage	X	Eu, As	A. Konstantinov, pers. comm.
		Galeruca	pomonae (Scopoli)	rosettes	X	Но	Doguet (1994); Steinhausen (1996)
		Galeruca	tanaceti (L.)	Foliage	X	Eu	A. Konstantinov, pers. comm.
		Longitarsus	brisouti Heikertinger	Foliage	X	Eu	Doguet (1994); Biondi (1996)
		Longitarsus	longipennis Kutschera	Roots, foliage	X	Eu, As	Biondi (1996); Cagán et al (2000)
		Longitarsus	luridus (Scopoli)	Bolting plants	X	Pa	Doguet (1994); Biondi (1996)
		Longitarsus	pratensis (Panzer)	Foliage	X	Med	Doguet (1994); Biondi (1996)
		Longitarsus	strigicollis Wollaston	Bolting plants, rosettes	G	Med	Doguet (1994); Biondi (1996)
		Neocrepidodera	ferruginea(Scopoli)	Foliage	X	Eu, As	Warchalowski (2003)
		Phyllotreta	nigripes (Fab.)	Roots, flowers	X	Med	Doguet (1994); Warchalowski (2003)
		Psyllioides	circumdatus (Redtenbacher)	Foliage	X	Med	Doguet (1994); A. Konstantinov, pers. comm.
		Sermylassa	halensis (L.)	Foliage	X	Eu, As	Warchalowski (2003)
		Smaragdina	limbata (Stéven)	Foliage	X	Med, As	Gök (2002); A. Konstantinov, pers. comm.
		Smaragdina	xanthaspis (Germar)	Foliage	X	Eu, AM	Warchalowski (2003)
Diptera	Agromyzidae	Chromatomyia	ramosa (Hendel)	Leaf miner,	G	Eu	Hering (1957)
Hemiptera	Coreidae	Coreus	marginatis (L.)	seeds?	X	Eu, As	Harizanova, unpublished data
	Miridae	Lygus	spp.	Foliage, heads	X	Co	Harizanova, unpublished data
Homoptera	Aphididae	Myzus	persicae (Sulzer)	Foliage	X	Co	Stoner (1951)
•	Cercopidae	Aphrophora	spp.	Rosettes	X	Co	Harizanova, Rector and Sforza,
		Cercopis	vulnerata Illiger	Foliage	X	Со	unpublished data Harizanova, Rector and Sforza, unpublished data
		Philaenus	spumarius (L.)	Foliage	X	На	Harizanova, Rector and Sforza,

Table 1 (continued)

Order	Family	Genus	Species (author)	Damage	Specificity ^a	Rangec	Reference(s)
Hymenoptera	Cimbicidae	Abia	sericea (L.)	Foliage	X	Eu, AM	André (1879); Taeger et al. (1998); Magis (2001); Liston (1995)
	Tenthredinidae	Macrophya	sp.	Bolting plants	U	Eu, AM	Vassilev (1978); D. Smith, pers. comm.
		Macrophya	diversipes (Schrank)	Unknown	X^b	Med, As	Çalmasur and Özbek, 2004
		Macrophya	postica (Brullé)	Unknown	\mathbf{U}^{b}	Eu, AM	D. Smith, pers. comm.
Lepidoptera	Adelidae	Nemophora	metallica (Poda)	Flowers, seeds	X	Eu	Novak et al. (1983)
1 1	Arctiidae	Diaphora	mendica (Clerck)	Foliage	X	Pa	Koch (1984); Dubatolov (1996)
		Rhyparia	purpurata (L.)	Foliage	X	Pa	Koch (1984); Dubatolov (1996)
	Geometridae	Synopsia	sociaria (Hübner)	Foliage, bracts	X	Eu, As	Koch (1984)
	Noctuidae	Acronicta	rumicis (L.)	Foliage	X	Eu, As	Novak et al. (1983); Koch (1984)
		Autographa	gamma (L.)	Foliage	X	Eu, As	M.G. Pogue, pers. comm.
		Heliothis	viriplaca (Hufnagel)	Foliage, seeds	X	Eu, As	Novak et al. (1983); Koch (1984)
		Tyta	luctuosa (Den. and Schiff)	foliage	X	Eu	Koch (1984); Rosenthal et al. (1988)
		Xestia	c-nigrum (L.)	Foliage	X	На	Koch (1984); Ferguson et al. (1999)
	Nymphalidae	Euphydryas	aurinia (Rottemburg)	Rosettes, bolting plants	X	Eu, AM	Mazel (1986); Wahlberg (2001)
	Tortricidae	Cochylis	roseana (Haworth)	Seeds	X	Eu, Med	Cheesman (1996); O. Cheesman, pers. comm.
		Endothenia	gentianaeana (Hübner)	head cavity	X	Eu, As	Gibeaux (1988); Cheesman (1996); Trematerra and Baldizzone (2004)
		Endothenia	oblongana (Haworth)	Root crown, stalk	X	Eu	Gibeaux (1988); Trematerra and Baldizzone (2004)
Viruses Virus	Unknown	Unknown	unknown	Symptoms at bolting	U	Eu	Rector and Widmer, unpublished data (see text)

^a Specificity index: X, not specific to Dipsacaceae; G, only known from Dipsacaceae, including the genus *Dipsacus*; S, only known from *Dipsacus* spp.; and U, unknown.

include host specificity testing for potential non-target plants in the proposed region of release followed by mass-rearing and redistribution of the BCC in the release area, assuming favorable host specificity test results. This would be similar to a classical augmentation biological control strategy (Pedigo, 1989) unless this moth currently shows only a restricted distribution in the USA with respect to the target. There may also be Old World insects that have been introduced into N. America that feed on teasels. The polyphagous European chrysomelid flea beetle Longitarsus luridus (Scopoli), which has been collected by the authors in the adult stage feeding on teasel foliage in Bulgaria and Romania, has been reported by Doguet (1994) as accidentally introduced into N. America. There are, however, no reports of it feeding on teasels there. Although L. luridus itself is not a very suitable BCC due to its highly polyphagous habit, there may be other accidentally introduced teasel-feeding insects that could be more interesting. Indeed, precedents for such serendipidous biological control agents exist (e.g., McClay, 1990).

In addition to these herbivores, Stoner (1951) described a virus attacking cultivated teasel (*D. sativus*) rosettes in central California. He found that it was nonpersistently transmitted by both *Myzus persicae* (Sulzer) and *Macrosiphum rosae* (L.) and in a test of several horticulturally important plant species, the virus would only attack teasel and *Scabiosa atropurpurea* L., an ornamental Dipsacaceae that has been introduced to the US (Bailey, 1951). A survey of the area near this virus's discovery, and perhaps an intentional planting of one or more small fields of teasels in an attempt to "trap" this virus could be worthwhile. Additional general surveys for herbivores in the invaded range are needed to further assess the possible presence of BCCs that may already exist in N. America.

^b Species possibly accidental; not found feeding on *Dipsacus*.

c AM, Asia minor; As, Asia; Co, cosmoplitan; Eu, Europe; Ha, Holarctic; Med, Mediterranean; NA, North America; Pa, Palearctic; and U, unknown.

Table 2 Invertebrate herbivores and pathogens associated with plant species in the family Dipsacaceae that have not been collected by the authors

Order	Family	Genus	Species (author)	Damage	Specificity ^a	Range ^b	Reference(s)
Fungi							
Agnomycetales	Agnomycetaceae	Sclerotium	rolfsii Sacc.	Root, stem	X	NA	Farr et al. (2004)
Dothideales	Dothideaceae	Didymella	exigua (Niessl) Sacc.	Stem, foliage	X	Co	Farr et al. (2004)
		Ramularia	silvestris Sacc.	Foliage	S	NA	Farr et al. (2004)
	Hysteriaceae	Clathrospora	permunda (Cooke) Sacc.	Foliage	X	Но	Farr et al. (2004)
Erysiphales	Erysiphaceae	Erysiphe	cichoracearum DC	Stem, foliage	X	Co	Farr et al. (2004)
		Erysiphe	communis (Wallr.) Link	Stem, foliage	X	Eu	Farr et al. (2004)
		Erysiphe	galeopsidis DC	Stem, foliage	X	As	Farr et al. (2004)
		Erysiphe	knautiae Duby	Stem, foliage	G	Eu	Farr et al. (2004)
		Phyllactinia	guttata (Wallr.) Lév.	Foliage	X	Но	Farr et al. (2004)
		Sphaerotheca	dipsacearum (Tul. and C. Tul.)	Foliage	G	Eu	Farr et al. (2004)
		Sphaerotheca	fuliginea (Schlect.) Pollacci	Foliage	X	Co	Farr et al. (2004)
Melanommatales	Didymosphaeriaceae	Diapleella	clivensis (Berk. & Broome)	Foliage	X	NA	Farr et al. (2004)
Moniliales	Dematiaceae	Cercospora	elongata Peck	Foliage	S	As, NA	Farr et al. (2004)
		Fusariella	hughesii ChabFrydm	Foliage	X	Co	Farr et al. (2004)
	Moniliaceae		omnivorum Duggar	Foliage	X	NA	Farr et al. (2004)
Myaaanhaarallalaa	Mycosphaerellaceae	Sphaerella	asterinoides Ellis & Everh.	Stem, foliage	S	Eu, NA	Farr et al. (2004)
		•					
Peronosporales	Peronosporaceae	Peronospora	dipsaci Tul. ex de Bary	Stem, foliage	G		Farr et al. (2004)
Pleosporales	Pleosporaceae	Leptosphaeria	conoidea (De Not.) Sacc.	Root, stem	X	NA	Farr et al. (2004)
		Pleospora	herbarum (Fr.) Rabenh.	Foliage	X	Eu, As	Farr et al. (2004)
		Pleospora	scrophulariae (Desm.) Höhn	Foliage	X	As, NA	Farr et al. (2004)
		Pleospora	vulgaris Niessl	Foliage	X	As	Farr et al. (2004)
	Venturiaceae	Venturia	cephalariae Kalchbr.& Cooke	Foliage	G	As	Farr et al. (2004)
Sphaeropsidales	Sphaeropsidaceae	Ascochyta	dipsaci Bubák	foliage	G	Eu	Farr et al. (2004)
		Phoma	oleracea var. dipsaci	Crown, root,	S	NA	Farr et al. (2004)
			Sacc.	foliage			· · ·
Mites				-	_	_	
Acarina	Eriophyidae	Aceria	squalida (Nat.)	Flowers	F	Eu	Petanovic and Stankovic (1999)
		Epitrimerus	knautiae Liro	Flower galls?	G	Eu	Petanovic (2001)
Insects							
Coleoptera	Buprestidae	Trachys	puncticollis Obenberger	Leaf miner	X	Eu	Curletti (1994)
		Trachys	troglodytes Gyllenhal	Leaf miner	F	Eu	Hering (1957)
	Chrysomelidae	Longitarsus	allotrophus Furth	Foliage	X	Eu	Doguet (1994);
	•		•	•			Biondi (1996)
Diptera	Agromyzidae	Agromyza	dipsaci Hendel	Leaf miner, bolting plants	S	Eu	Hering (1957)
		Agromyza	woerzi Groschke	Leaf miner	F	Eu	Hering (1957)
		Aulagromyza	similis (Brischke)	Leaf miner	X	Eu	Hering (1957)
		Chromatomyia	horticola (Goureau)	leaf miner	X	Co	Hering (1957)
		Chromatomyia	scabiosae (Hendel)	Leaf miner	F	Eu	Hering (1957)
		Chromatomyia		Leaf miner	F		
		•	scabiosarum (de Meijere)			Eu	Hering (1957)
		Chromatomyia	succisae (Hering)	Lleaf miner	G	Eu	Hering (1957)
		Liriomyza	strigata (Meigen)	Leaf miner	X	Eu	Hering (1957)
		Melanagromyza	knautiae Hering	leaf miner	F	Eu	Hering (1957)
Homoptera	Aphididae	Aphis	chloris Koch	Foliage	X	Eu	Patch (1938); Briese (1989)
		Aphis	confusa Walker	foliage	F	Eu	Patch (1938)
		Aphis	conspersa Walker	Foliage	F	U	Patch (1938)
		Aphis	ochropus Koch	Foliage	X	Eu	Patch (1938)
		Aphis	rumicis L.	Foliage	X	Co	Patch (1938)
		Aphis	scabiosae Schrank	foliage	X	As	Patch (1938)
		Macrosiphum	rosae (L.)	Foliage	X	Co	Patch (1938)
		Macrosiphum	rudbeckiae Fitch	Foliage	X	NA	Patch (1938)
Hymenonters	Cimbicidae	•		-	X	Eu	
Hymenoptera	Cimbicidae	Abia	aenea (Klug)	Foliage	Λ	Eu	Taeger et al. (1998); Magis
		Abia	a an dana V	falia a-	E	E.,	(2001)
		Abia	candens Konow	foliage	F	Eu	Liston (1995); Magis (2001)

Table 2 (continued)

Order	Family	Genus	Species (author)	Damage	Specificity ^a	Range ^b	Reference(s)
		Abia	fasciata (L.)	Foliage	X	Eu	Magis (2001)
		Abia	lonicerae (L.)	Foliage	X	Eu	Chevin (2001);
				-			Magis (2001)
	Tenthredinidae	Macrophya	albicincta (Schrank)	Foliage	X	Med, As	Magis (2002);
		• •	, , ,	-			Çalmasur and
							Özbek, 2004
		Macrophya	crassula (Klug)	Foliage	X	Eu, AM	Magis (2002);
			-	-			Çalmasur and
							Özbek (2004)
		Tenthredo	atra L.	Foliage	X	Eu, As	André (1879);
							Lacourt (1999)
Lepidoptera	Arctiidae	Artimelia	latreillei (Godart)	Foliage	X	Eu	de Freina and
							Witt (1987)
		Ocnogyna	parasita(Hübner)	foliage	X	Eu, As	de Freina and
							Witt (1987)
	Coleophoridae	Coleophora	conspicuella Zeller	Foliage	X	Eu, As	Hering (1957)
		Coleophora	paripennella Zeller	Foliage	X	Eu	Hering (1957)
	Gracillariidae	Phyllonorycter	scabiosella (Douglas)	Leaf miner	F	Eu	Emmet et al.
							(1985)
	Noctuidae	Eriopygodes	imbecilla (Fab.)	Foliage	X	Eu, As	Novak et al.
							(1983)
		Nola	chlamitulalis (Hübner)	foliage	X	Med, As	de Freina and
							Witt (1987)
		Papaipema	arctivorens Hampson	Root	X	NA	Hodges (1983);
							Covell (1984)
	Nymphalidae	Euphydryas	desfontainii (Godart)	Rosettes, bolting	G	Eu	Nóvoa Pérez
				plants			and García-
							Villanueva
							(1996);
							Wahlberg (2001
	Oecophoridae	Agonopterix	arenella (Den. and Schiff.)	foliage	X	Med, NA	Novak et al.
							(1983); Harper
							et al. (2002)
		Agonopterix	kaekeritziana (L.)	Foliage	X	Eu	Novak et al.
							(1983)
	Papillionidae	Parnassius	apollonius (Eversman)	Foliage	X	As	Novak et al.
						_	(1983)
	Psychidae	Apterona	helicoidella (Vallot)	Foliage	X	Eu	Hering (1957);
							Novak et al.
				- ·	_	_	(1983)
	Pterophoridae	Gillmeria	miantodactylus Zeller	Foliage, flowers	F	Eu	Gielis (1996)
		Stenoptilia	annadactyla Sutter	Foliage, flowers	F	Eu	Gielis (1996)
		Stenoptilia	aridus (Zeller)	Foliage, flowers	X	Med	Gielis (1996)
		Stenoptilia	bipunctidactyla (Scopoli)	Foliage, flowers	X	Eu	Gielis (1996)
	0 4 111	Stenoptilia	stigmatodactylus (Zeller)	Foliage, flowers	X	Med	Gielis (1996)
	Scythrididae	Scythris	picaepennis (Haworth)	leaf miner	X	Eu, As	Hering (1957);
							Bengtsson
		G .1 :	: 11 (7.11)	т. с. :	37	Г	(1997)
	0.1: :1	Scythris	siccella (Zeller)	Leaf miner	X	Eu	Hering (1957)
	Sphingidae	Acherontia	athropos (L.)	Foliage	X	Af, Eu	Pittaway (1993)
		Hemaris	croatica (Esper)	Foliage	X	Eu, AM	Pittaway (1993)
		Hemaris	fuciformis (L.)	Foliage	X	Eu, As	Pittaway (1993)
		Hemaris	tityus (L.)	Foliage	X	Eu, As	Pittaway (1993)
	Tortricidae	Hyles	livornica (Esper) hartmanniana (Clerck)	Foliage	X F	Af, Eu, As	Pittaway (1993)
	Tortricidae	Aethes	narimanniana (Cierck)	Root, crown	Г	Eu	Razowski
							(1970);
							Trematerra and
							Baldizzone
		C	:	E-fina	v	Е.,	(2004)
		Cnephasia	incertana (Treitscke)	Foliage	X	Eu	Hering (1957)
		Cnephasia	stephensiana (Doubleday)	Foliage	X	Eu Mad	Hering (1957)
		Cochylimorpha	straminea (Haworth)	Flower buds, seeds	Λ	Eu, Med	Novak et al.
		Cochylis	flaviciliana (Westwood)	Flower buds, seeds	C	Eu, Med	(1983) Razowski (1970

(continued on next page)

Table 2 (continued)

Order	Family	Genus	Species (author)	Damage	Specificitya	Range ^b	Reference(s)
		Diceratura	ostrinana (Guenée)	Immature flowers	X	Eu, AM	Razowski (1970); Gibeaux (1988)
		Endothenia	marginana (Haworth)	Head cavity, seeds	X	Eu	Gibeaux (1988); Trematerra and Baldizzone (2004)
		Rhopoboda	stagnana (Den. and Schiff.)	Leaf miner	F	Eu	Hering (1957)
Nematodes Tylenchida	Anguinidae	Ditylenchus	dipsaci Filipjev	Roots	X	На	Thorne (1945)
Viruses Virus	Potyviridae?	??	??	Symptoms in rosettes	G	NA	Stoner (1951)

^a Specificity index: X, not specific to Dipsacaeea; F, only known from the family Dipsacaeeae but not known from *Dipsacus*; G, only known from Dipsacaeeae, including the genus *Dipsacus*; S, only known from *Dipsacus* spp.; and U, unknown.

4.3. Candidates from the native range of D. fullonum and D. laciniatus

Despite the long history of teasel cultivation in Europe and the United States, there is very little reference in the scientific literature to teasel pests, beside the virus described by Stoner (1951) and a reference to the "teasel fly," Chromatomyia ramosa (Hendel) (Diptera: Agromyzidae), which was considered an important enough pest in England to merit regularly scheduled sprays of malathion or DDT (Topham, 1968). This insect was reared from mines in D. fullonum rosette leaves from France and Greece in the spring of 2004. The insect is bivoltine with a larval generation from May to August and another from October to April and it is also reported to feed on new growth at the center of the rosette late in the larval stage (Hering, 1957). This insect's host range is limited to the family Dipsacaceae (Hering, 1957) and three congeners: C. scabiosae (Hendel); C. scabiosarum (de Meijere); and C. succisae (Hering) are also only known to feed on Dipsacaceae (Hering, 1957). Another agromyzid leaf mining fly, Agromyza dipsaci Hendel, has also only been described feeding on D. fullonum (Hering, 1957). D. fullonum leaves collected in France in the summer of 2004 were observed to have damage similar to that of A. dipsaci as illustrated by Hering (1957), although it was too late in the season to collect insects from these leaves. This species is univoltine with the larval feeding occurring in May and June along the margins of the stem leaves of bolted plants.

A tortricid seed-feeder, *Cochylis roseana* (Haworth), has been well-documented from *D. fullonum* (Cheesman, 1996) and larvae matching this species's description have been collected from seedheads of *D. laciniatus* in Bulgaria. Female moths lay clusters of eggs at the bases of involucral bracts on young teasel flowerheads, whereupon hatched larvae enter the head and begin feeding on florets and developing seeds before switching exclusively to seeds (Cheesman, 1996). Larvae of this species feed gregariously and Cheesman (1996) reported an average of more than 18

larvae per head. They overwinter in the seedhead as larvae before pupating in the spring and emerging as adults shortly thereafter. This species is also known to attack *Solidago* spp., a group outside the order Dipsacales of which there are several N. American species. This fact would complicate host-specificity testing on *C. roseana*. A congener, *C. flaviciliana* (Westwood) could be a more promising candidate. It has been reported from teasel flowerheads and its host-range is only known to include plants in the Dipsacaceae (Table 2). However, *C. flaviciliana* has not yet been collected by the authors. Another congener (*C. hospes* Walsingham) is a known pest of sunflower (Barker, 1996), while another [*C. atricapitana* (Stephens)] has been released as a biological control agent against *Senecio jacobaeae* L. in Australia (McLaren, 1992).

Larvae of two species in the nymphalid genus Euphydryas Scudder, E. aurinia (Rottemburg) and E. desfontainii (Godart), are known to feed on *Dipsacus* spp. (Mazel, 1986; Wahlberg, 2001). E. aurinia has been encountered in the field in several countries across Europe from France to Turkey (Sforza, 2004). However, this gregarious feeder also feeds on host plants in at least three families, including one (Gentianaceae) outside the order Dipsacales (Wahlberg, 2001). The native range of E. desfontainii covers most of the Iberian peninsula, parts of N. Africa and isolated populations in SW France, where it is a protected species. Its hostrange appears to be narrower than that of E. aurinia, feeding only on plants in the Dipsacaceae (Wahlberg, 2001). Females of both of these species lay groups of eggs on the host plant in May and June. The larvae hatch soon after and feed briefly, completing early instars before aestivating until autumn. The larvae may feed again in autumn before hibernating until early spring when they complete feeding, pupate, and emerge as adults.

Longitarsus strigicollis Wollaston is a chrysomelid flea beetle that has been found feeding in large numbers on D. fullonum in France and Italy. This species is found in the adult stage whenever the weather is warm enough and it is thought for this reason to be multivoltine in warmer

b Af, Africa; AM, Asia Minor; As, Asia; Co, cosmoplitan; Eu, Europe; Ha, Holarctic; Med, Mediterranean; NA, North America; and U, unknown.

climates (Doguet, 1994). The adults of this species are only known to feed on hosts in the Dipsacaceae (Doguet, 1994) but the larval stage is as yet undescribed. Based on information about the larval habits of congeneric species (Steinhausen, 1996), it is believed the larvae of *L. strigicollis* may be external root feeders of Dipsacaceae, overwintering in the soil as immatures. This hypothesis is currently being tested.

The cerambycid stalk borer *Agapanthia osmanlis* Reiche feeds only on hosts in the Dipsacaceae (Kovacs, 1998; Rejzek et al., 2003) and has been collected on *Dipsacus* spp. in Bulgaria and Turkey. Adults of this species are active in the spring and summer and feed on the foliage of bolting plants. The female lays single eggs inside teasel stalks where the larva feeds internally. After overwintering, the insect pupates in the spring before emerging as an adult in summer. Tests are underway to determine whether the larval feeding causes damage early enough in the plant's life to significantly affect seed production.

Larvae of the cimbicid sawfly *Abia sericea* (L.) were found feeding on *D. laciniatus* in Bulgaria and Turkey. This species has been described from several hosts in the Dipsacaceae but also from *Fragaria* sp. (strawberries) (Liston, 1995). Tests are currently underway to investigate the validity of this report, since it would represent the only known host record for a European *Abia* sp. from outside the Dipsacaceae or the closely related Caprifoliaceae (Taeger et al., 1998). Indeed, as Taeger et al. (1998) state explicitly, "under field conditions *Fragaria* is surely not a host plant (of *A. sericea*)." The congener *A. candens* Konow, is only known from Dipsacaceae (see Table 2).

The eriophyid mite *Epitrimerus knautiae* Liro has been collected from flower galls of *D. laciniatus* in Yugoslavia (Petanovic, 1999; R. Petanovic, pers. comm.). It is not clear whether these mites caused the galls within which they were found (R. Petanovic, pers. comm.). The only other reports of this mite list it as vagrant on leaves of *Knautia arvensis* (L.) Coulter, a close relative of *Dipsacus*, in Finland (Liro, 1942) and Poland (Boczek, 1964).

Several fungi have been collected from *Dipsacus* spp. and shown to be primary pathogens of *D. fullonum* (see Table 1). Among these, *Sclerotinium sclerotiorum* (Lib.) de Bary may hold promise as a mycoherbicide, although it has a broad host range. Attenuated mutants of this species have been proposed for control of *Cirsium arvense* (L.) Scop. in the US (Brosten and Sands, 1986) and *Chrysanthemoides monilifera* (L.) T.Nord in Australia (Cother et al., 1996). Depending on the success of *S. sclerotiorum* in being approved for release and in controlling these or other targets, *D. fullonum* and *D. laciniatus* could become new targets for this mycoherbicide.

In addition to the three fungi that the authors have collected on *Dipsacus* in the field (Table 1), reports of 24 other fungi associated with *Dipsacus* spp. are cataloged in the USDA-ARS GRIN database (Farr et al., 2004) (Table 2). Although some genera of these fungi are known to have broad host ranges, individual isolates may show higher

specificity. Thus, it is important to isolate fungi from teasels and conduct further tests from these specific isolates on a case-by-case basis. As we found little work has been done on any of the fungi listed in Table 2, it cannot be said with certainty what potential any of these fungi have as BCCs. Exploration and research remains to be conducted to collect and evaluate individual fungal isolates for host specificity and impact.

In addition to the four fungal species listed in Table 2 as specific to *Dipsacus* (all of which are based on only a few reports per fungal species), two powdery mildews (family Erysiphaceae), have potential as promising BCCs from the standpoint of specificity, although their impact on the plant is not yet known. *Sphaerotheca dipsacearum* (Tul and C. Tul) has 58 host records in the GRIN database from seven host genera, all of which are in the Dipsacaceae (Farr et al., 2004). For *Erysiphe knautiae* Duby, only two of the 221 host records in the GRIN database fall outside the Dipsacaceae (*Papaver anomalum* Fedde. and *Saxifraga manchuriensis* (Engler) Komarov) (Farr et al., 2004). These plant species would need to be tested in any subsequent host specificity testing.

Symptoms of an as yet unidentified virus have been observed on *D. fullonum* in southern France and similar symptoms have been observed on *D. fullonum* and *D. laciniatus* in Bulgaria, northern France, Slovenia, and Turkey. This virus causes mosaic-like, chlorotic symptoms in bolting plants, followed by stunting and discoloration of the mature plant. These symptoms are quite unlike those of the teasel virus described by Stoner (1951). Field observations of tagged plants with viral symptoms indicate a significant reduction in number of seedheads per plant, as well as a proportion of plants that produce no seed at all (Rector, unpublished data). Work is under way to identify this virus and to determine how it is vectored from one generation to the next.

5. Risk assessment and potential interactions of biological control candidates

As BCCs are prioritized for further work, host-specificity test plant lists need to be agreed upon. As agents with narrow host ranges tend to be restricted to related host plants (Wapshere, 1974), phylogenetic clade information to develop these lists from recent molecular phylogenies should be used (Briese, 2002). For Dipsacales, three recent molecular phylogenetic treatments (Bell et al., 2001; Donoghue et al., 2001; Zhang et al., 2003) are all in general agreement and should assist this.

Potential interactions between the aforementioned highpriority BCCs have not yet been evaluated. Any competitive or synergistic interactions between multiple BCCs approved for release can be tested before their respective introductions since no teasel biological control agents have yet been released and there currently appears to be little natural enemy activity on teasel in N. America. Priority for study of subsequent BCCs should however be given to candidates attacking different plant parts from the first released agents, as advocated by Malecki et al. (1993) and others.

6. Conclusions

As invasive teasels continue to spread in the US, particularly on lands that are not intensively managed for weed control, the need for a self-sustaining management strategy, such as biological control, increases. Teasels present particular opportunities as biological control targets, given the absence of any North American relatives or economically important plants within the family. In their native ranges, D. fullonum and D. laciniatus rarely achieve the population sizes or densities that have induced five American states to list either or both as noxious species. Whether natural enemies of these teasels are responsible for keeping native populations in check is not known, however based on the results of the initial literature and field surveys presented in this article, it appears that natural enemies of Dipsacus spp. are both numerous and specific enough to yield promising biological control agents that could suppress invasive populations.

Among the BCCs collected and identified to date, the highest priority for initial study has been assigned to two insects that attack the first-year vegetative rosette stage of teasel. Damage at this stage appears to be the most promising for biological control since it could delay flowering from one summer to the next and perhaps prevent it altogether. These two insect species are the chrysomelid flea beetle L. strigicollis, which feeds on foliage as an adult and may also feed on roots in the larval stage and the agromyzid fly C. ramosa, which mines rosette leaves and may feed at the apical meristem late in larval development. Among those BCCs identified in the literature but not yet observed or collected by the authors in the field, those to which the authors assign the highest priority for targeted surveys include the foliage-feeding nymphalid butterfly Euphydryas desfontainii, the eriophyid mite E. knautiae, the root-boring noctuid moth P. arctivorens, an aphid-transmitted virus described from California (Stoner, 1951), and the powdery mildews *E. knautiae* and *S. dipsacearum*.

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